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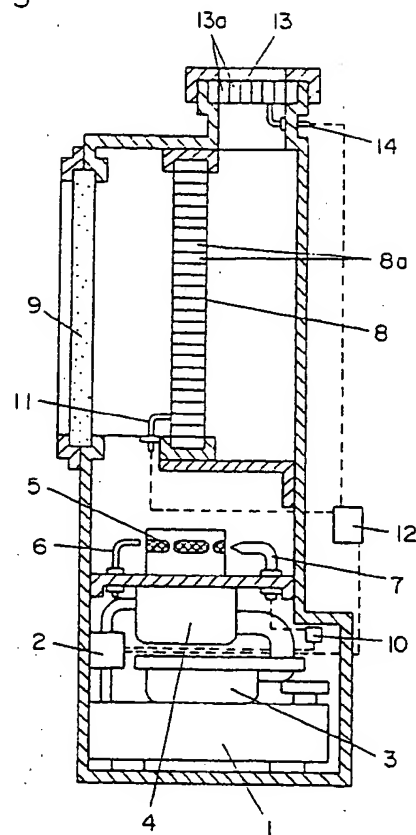
(54) CATALYTIC COMBUSTION APPARATUS.

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 (57) A catalytic combustion apparatus in which a flame port (5) equipped with an ignition electrode (6) and a flame rod (7) nearby thereof, is arranged on the downstream of a mixing chamber (4) where the fuel and the air are mixed together, a catalyst layer (8) having many communication holes (8a) is provided on the downstream side thereof, ignition means (6) is operated to form flame at the flame port (5), supply of the fuel is once stopped after a

predetermined period of time has passed to extinguish the flame, and the fuel is supplied again without operating the ignition means (6) such that the combustion reaction takes place on the surface of the catalyst layer (8). When the flame is formed at the flame port (5), it is detected that a predetermined current is not obtained from said ionic current detect means (7). When the combustion reaction is started on the catalyst layer (8), on the other hand, the

current that is obtained is detected to stop the combustion.

FIG. 3



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DESCRIPTION

1 TITLE OF INVENTION

CATALYTIC BURNING APPARATUS

TECHNICAL FIELD

The present invention relates to a catalytic
5 burning apparatus for effecting an oxidizing reaction of
fuel on a solid oxidizing catalyst.

BACKGROUND ART

Heretofore, several apparatus for effecting an
oxidizing reaction of liquid or gaseous fuel on a solid
10 oxidizing catalyst have been proposed, for example, an
apparatus as shown in Fig. 1 (Catalyst, Vol. 29, No. 4,
313, 1987).

In Fig. 1, numeral 101 denotes a fuel pipe,
numeral 102 ejection ports, numeral 103 an insulator layer,
15 numeral 104 an electric heater, numeral 105 a catalyst
layer, and numeral 106 a cover. Fuel is supplied through
the ejecting ports 102 formed in the fuel tube 101 in a
distributed manner, and passed through the porous insulator
layer 103 to the catalyst layer 105 which is preheated by
20 the electric heater 104. On the other hand, air is supplied
from the underside of the cover 106 under the function of
convection. Near the surface of the catalyst layer 105,
the fuel and the air are mixed with each other by diffusion,
and a catalytic burning is effected on the fibered porous

1 catalyst layer 105.

The catalytic burning apparatus of this type, however, has problems as follows. Firstly, it is required to heat the catalyst layer 105 to a temperature at which 5 the catalytic reaction starts, and it takes a long time to heat the catalyst layer to the predetermined temperature by the electric heater 104, unless a heater of a great capacity is used. Secondly, since the catalyst layer 105, from the surface of which the heat 10 is radiated forwards, is only covered in a halfly exposed manner by the cover 106 made of such as a porous metal, there is a fear that the burning is interrupted by a gust or a water spray, frequently causing an imperfect combustion and producing an offensive smell and a harmfull 15 carbon monoxide. Thirdly, when the apparatus is used for a long time and the activity of the catalyst layer is deteriorated, there occurs a fear that the imperfectly burned fuel flows out, and an offensive smell and a great amount of harmful carbon monoxide are continuously 20 produced due to the imperfect combustion, because there is provided no detecting means for detecting the deterioration of the catalyst layer. Fourthly, in the case where the fuel is burned in a closed space such as in a room, the burning is not stopped as far as the 25 temperature of the catalyst layer is maintained in a predetermined range, even when the oxygen density has been decreased to a level having an adverse influence on the human health, thereby causing a continuation of the oxygen

1 starvation and the imperfect combustion.

DISCLOSURE OF INVENTION

The present invention provides a catalytic burning apparatus which can solve the above-mentioned problems and
5 is superior in burning control capability and in safety.
The present invention has a characterizing feature that flame ports added with ignition means and ion current detecting means are disposed upstream of the catalyst layer, and an abnormal combustion environment or combustion
10 condition is detected based on the ion current value.

BRIEF DESCRIPTION OF DRAWINGS

Fig. 1 is a structural view of a catalytic burning apparatus of a prior art,

Fig. 2 is a structural view of a catalytic
15 burning apparatus according to a first embodiment of the present invention,

Figs. 3, 4, 5 and 6 are structural views of catalytic burning apparatus according to second, third, fourth and fifth embodiments of the present invention,
20 respectively,

Fig. 7 is a performance illustration for showing variation of transforming rates in oxidizing reaction on kerosine or carbon monoxide due to the composition of precious metals,

25 Fig. 8 is a performance illustration for showing an influence of the ratio of the auxiliary catalyst volume

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1 to the catalyst layer volume on the transforming rates
in oxidizing reaction on kerosine or carbon monoxide,
and

Fig. 9 is a performance illustration for showing
5 an influence 5 of the cell number of auxiliary catalyst
layer on the transofrming rate in oxidizing reaction or
the carbon monoxide.

BEST MODE FOR CARRYING OUT THE INVENTION

Embodiments of the present invention will be
10 described below. Figs. 2 to 6 relate to embodiment of
the present invention, and in these figures, the same
constituent members are indicated with the same numerals.
Figs. 7 to 9 relate to catalytic performances showing
influences of the structure of catalyst layer or
15 auxiliary calsyts layer and composition of the precious
metals on the oxidizing reaction or kerosine or carbon
monoxidide.

In Fig. 2, numeral 1 denotes a liquid fuel
tank, numeral 2 a fuel pump, numeral 3 an air blast
20 fan, numeral 4 a mixing room. At the exit of the mixing
room 4 are provided flame ports 5, and near the flame ports
5 are provided an ignition plug 6 and an electrode for
measuring the ion current in the flame, i.e. so-called
a flame rod 7.

25 Above the flame ports 5 is provided a vertically
arranged catalyst layer 8 which includes an active compo-
sition of platinum metal carried out a honeycome-like

1 ceramic flat plate mainly composed of silica-alumina
and bored with a plurality of communicating holes 8a.
Upstream of the catalyst layer 8 (front side) is arranged
a transparent window 9 made of a glass plate and located
5 opposite to the catalyst layer 8. Numeral 10 denotes a
control section for the pump 2, numeral 11 a thermocouple
for detecting the temperature of the catalyst layer 8,
and numeral 12 a burning control circuit.

Next, the operation will be described in
10 detail. The fuel (kerosine) supplied from the fuel pump
2 is vaporized in the mixing room 4, sufficiently
premixed with the air supplied from the fan 3, and
transferred to the flame ports 5 locating above. Firstly,
the mixed gas is ignited at the flame ports 5 by the
15 ignition plug 6, thereby starting a flame burning. The
exhaust gas of high temperature flows upwards passes
through the communicating holes 8a and flows to downstream
side, while the temperature of the catalyst layer is
raised. When, after burning for a predetermined time
20 length, the thermocouple 11 detects that the temperature
of the catalyst layer 8 reaches a sufficiently high
temperature, the pump 2 is once stopped for putting out
the flame, and is started again. In this process, the
premixed gas coming from the mixing room 4 flows to the
25 catalyst layer 8 which is vertically arranged above.
Since the catalyst layer 8 has been sufficiently heated,
the mixed gas effects catalytic burning mainly at the
upstream side (front surface) surface, and the burned

1 exhaust gas flows to the downstream side (rear surface)
through the communicating holes 8a. A part of the reaction
heat generated at the surface of the catalyst layer 8
penetrates through the transparent window 8, and another
5 part of the reaction heat heats the transparent window
8 and is radiated from the window as a secondary radiation,
these heats being radiated to the front side and used
for room heating or the like. At the ignition time when
the flame is formed at the flame ports 5, the flame rod 7
10 confirms that an ion current of a predetermined flow rate
is flowing in the flame, and whereby a misignition or a
misfire is detected.

On the other hand, at the time when the flame
at the flame ports 5 has been extinguished and the
15 catalytic burning on the catalyst layer 8 has been
started, the flame rod 7 confirms, in contrast with the
above, that no flame exists at the flame ports 5, in
other words, no ion current is flowing, thereby detecting
that the burning has been completely switched into the
20 catalytic burning, and any flame due to an incomplete
extinguishment or a back-fire from the catalyst layer 8 to
the flame ports 5 does not exist at the flame ports 5.

By utilizing the flame heat produced at the
flame ports 5 for preheating the catalyst layer 8, the
25 whole amount of the high temperature exhaust gas is
passed through the communicating holes 8a of the catalyst
layer 8, thereby uniformly heating the whole region of
the catalyst layer 8. As a result, an efficient preheating

1 can be achieved. For example, the time required for pre-
heating the catalyst layer 8 to a predetermined temperature
is about 3 to 5 minutes in case of using an electric
heater of 1.5 kW, while it is not more than one minute in
5 case of using a flame burning of 1200 kcal/h. Further,
in case of an electric heater, the temperature is easily
raised near the heater, but very slowly raised at
the region remote from the heater, while in case of a
flame burning, the temperature is uniformly raised in a
10 short time without any local unevenness of the temperature.
In addition, there is not any fear that an electric
heater suffers an oxidizing corrosion or a heat damage
near the catalyst layer 8 which is constantly under high
temperature and oxidizing condition. Further, since an
15 abnormality in a burning start or in a catalytic burning
is always detected by the flame rod 7, a favorable
result can be obtained with respect to life length or
stability and safety of burning.

Although, in the above-mentioned arrangement,
20 the combustion air is totally supplied to the mixing
room 4, it is also possible to supply a part of the air
to near the flame ports 5 for effecting a diffusion
flame burning of the partially premixed gas. In this
case, the variation of the ion current is significant,
25 thereby improving the detecting precision of the flame rod
7 and assuring a surer detection of the flame burning
without deteriorating the perfect combustion feature of
the catalyst layer 8. The time length of the flame

1 burning required for preheating the catalyst layer 8
can be controlled by presetting it to a predetermined
value which is large enough for sufficiently raising the
temperature of the whole catalyst layer 8. However, it
5 is surer to detect the temperature of the catalyst layer
8 by means of a thermocouple 11 and confirm the tempera-
ture state. In the latter arrangement, in case of
a re-igniting just after a five extinguishment, where
the temperature of the catalyst layer is comparatively
10 high, there is obtained an advantage that an excessive
preheating can be omitted and a quick switching into a
catalytic burning can be carried out.

Further, the thermocouple 11 provided at the
catalyst layer 8 for detecting the preheating temperature
15 as mentioned above can also achieve a temperature control
function for catalytic burning. For example, it is
possible to detect an abnormal burning based on a drop of
the temperature of the catalyst layer 8, when the activity
of the catalyst layer 8 has been deteriorated, or the
20 catalyst layer has been partly damaged and the reaction
has become imperfect. In detail, in case the catalytic
activity is deteriorated, the central position of the
catalytic burning shifts from the upstream side (front
side) of the catalyst layer 8 to the downstream side (rear
25 side), and there occurs a temperature distribution change
that the temperature at the upstream side is lowered,
and the temperature at the downstream side is raised,
or the temperature of the downstream exhaust gas is raised.

1 By comparing these temperature distribution change with a
relation between fuel supply rate and temperature
distribution which is precalculated and stored in the
control circuit 12, an abnormal burning can be surely
5 detected, and the burning can be stopped based on the
detected abnormality. In case of a partial damage of
the catalyst layer 8, the fuel flows as gathering to
the damaged portion, and the temperature of the catalyst
layer 8 is lowered, thereby making it possible to detect
10 the abnormality. On the other hand, in case the surface
temperature of the catalyst layer 8 become significantly
high due to an abnormality of the pump 2 or the fan 3,
the temperature change is detected by the thermocouple
11, and a suitable control action such as indicating
15 an abnormality sign or stopping the burning can be
carried out, thereby assuring a safe and stable burning.

Although, in the above arrangement, a thermo-couple is used as temperature detecting means, any other temperature detecting means can be selected, for example,
20 a thermometer of a resistance type such as a thermistor or a thermometer of a radiation type using light. As to the location of the thermometer, it is not always necessary to locate the thermometer near the catalyst layer 8, but it is also possible to locate the thermometer in the exhaust gas passage as mentioned above for measuring the temperature of the exhaust gas, or to locate the same outside of the transparent window 9 for measuring the radiated heat amount. Since the catalyst layer 8 is located

1 in a closed passage extending downstream of the flame
ports 5, various external disturbing factors, for
example, a gust blowing in or a water spray, have no direct
influence on the catalyst layer 8 so that no imperfect
5 burning or no local misburning is caused, and a stable
and perfect burning can be maintained.

In case of kerosine catalytic burning having
an air ratio of about 1.5, the total amount of the oxygen
is sufficient, even if the oxygen density becomes as low
10 as 15%, in other words, the oxygen excessive ratio,
i.e. the ratio of an actual oxygen amount to a theoretically
required oxygen amount is maintained as high as about
1.1. In consequence, the burning reaction is maintained
at the catalyst layer 8. However, the oxygen density
15 in a room below 16% stands in an unsafe range having a
harmful influence on the human body. Here, during
catalytic burning, if a flame is formed at the flame
ports 5 by applying an electric current to the ignition
plug 6, and at the same time, the flame rod 7 is switched
20 to the flame detecting mode as seen in the preheating
process, an oxygen starvation state can be detected by
measuring the change of the ion current flowing through
the flame by means of the flame rod 7, because the state
of the flame and the ion density in the flame vary according
25 to the oxygen density. In case the ion current value
is beyond a predetermined value, an oxygen starvation is
concluded and the pump 2 is stopped through the controller
section 10 for interrupting the burning. Some flame ports

1 have a feature that, when the oxygen is starved, the
formation of a stable flame become difficult and the
flame blows out. In this case, the oxygen starvation
can be detected in a surer manner. By suitably setting
5 the electric current value, the burning can be stopped
when the oxygen density reaches 18% or 16%, thereby
preventing any unsafe operation. In this case, when the
ion current value is not beyond the predetermined value,
the fuel supply is temporarily interrupted similarly to
10 the ignition phase for extinguishing the flame at the
flame ports 5, and then the fuel supply is again started
for continuing the catalytic burning at the catalyst
layer 8. By conducting the above-mentioned operation for
a short time such as one to two minutes at intervals of
15 such as 30 minutes or one hour, the oxygen starvation
can be detected. further, since this operation is
controlled by the ignition plug 6 which is normally used
in the preheating process for the catalyst layer 8 and by
the flame rod 7 which is normally used for detecting
20 a misignition or a misfire, a sure safety can be assured
in a simple manner.

Next, a second embodiment will be described.
Referring to Figs. 3, downstream of the catalyst layer 8
is arranged an additional auxiliary catalyst layer 13,
25 which is also added with a thermocouple 14. The auxiliary
catalyst layer 13 is a honeycome-like ceramic plate
carrying an active composition of precious metals and bored
with a plurality of communicating holes 13a. Similarly to

1 the above-mentioned embodiment, a burning is started
through steps of forming a flame at the flame ports 5,
preheating the catalyst layer 8 and the auxiliary catalyst
layer 13 by using the combustion exhaust gas, extinguishing
5 the flame by once stopping the pump 2, and starting a
catalytic burning at the catalyst layer 8 by activating
the pump 2 again. The combustion exhaust gas further
flows upwards to the downstream side, and contacts with
the auxiliary catalyst layer 13, where the unburned fuel,
10 if any, is completely oxidized and thereafter exhausted
upwards through the communicating holes 13a as a clean
exhaust gas. In consequence, even when the fuel is not
completely burned at the catalyst layer 8 due to an
uneven preheating or an uneven temperature distribution,
15 the mixing is again effected and the mixed gas contacts
with the auxiliary catalyst layer 13 located downstream,
thereby completing the reaction and preventing any
unburned gas due to an imperfect combustion from being
exhausted. Further, even in case the activity of the
20 catalyst layer 8 has been deteriorated due to a long use,
the activity is compensated by the catalyst layer 13, and
a stable performance can be maintained for a long time.

In case the activity of the catalyst layer 8
drops down, the reaction position gradually shifts from
25 near the upstream side surface to the downstream side,
and finally, the fuel cannot be burned perfectly,
permitting a part of the fuel to pass therethrough in an
unburned condition or permitting carbon monoxide, which

1 is considered as an intermediate dissolved composition
or a reaction intermediate composition, to be mixed into
the exhaust gas. Accordingly, the temperature of the
catalyst layer 8 detected by the thermocouple 11 become
5 low. On the other hand, at the auxiliary catalyst layer
13 located at the downstream side, a combustion reaction
of the unburned fuel is effected, and due to this
reaction heat, the temperature of the auxiliary catalyst
layer 13 detected by a thermocouple 14 become high.
10 Thus, the temperature of the catalyst layer 8, which is
much higher than that of the auxiliary catalyst layer
13 at an initial stage, is gradually lowered relative
to the temperature of the auxiliary catalyst layer 13, and
finally the temperature relation between the two catalyst
15 layers is reversed. Even in this temperature reversed
condition, since a sufficient activity is maintained
at the catalyst layer 13, there is contained no unburned
fuel or carbon monoxide in the final exhaust gas, thereby
maintaining the exhaust gas at a clean state. Further,
20 in case the temperature difference between the temper-
atures detected by the thermocouple 11 and the thermocouple
14 become smaller than a predetermined value, this dif-
ference is judged to indicate a life limit of the catalyst
layer 8, and can be used as a signal for stopping the
25 burning. Thus, the deterioration of the catalyst layer
can be surely detected, and any imperfect combustion can
be prevented. The catalyst layer 8 may be arranged
vertically as shown in Fig. 3 and may be provided with a

1 transparent window at the upstream side for utilizing
the radiant heat, or may be, as seen in a third embodiment
shown in Fig. 4, provided with a air blowing fan 15 for
transforming the combustion heat into a warm wind for
5 room heating. Thus, there is no limitation with respect
to the arrangement of the catalyst layer 8 or to the
utilizing form of the reaction heat.

Next, a fourth embodiment will be described.
Referring to fig. 5, there is provided a secondary air
10 tube 16 which is branched from the outlet port of the
fan 3 and connected to a secondary air port 17 opening
at the upstream side of the auxiliary catalyst layer
13. Referring to an operational example where the
catalyst layer 8 and the auxiliary catalyst layer 13 are
15 preheated by burning the fuel at the flame ports 5, and
then the burning is switched to the kerosine catalytic
burning at the catalyst layer 8 with an air ratio 1.8 to
2.0, the surface temepratures of the catalyst layer 8
and the auxiliary catalyst layer 13 vary according to the
20 change of the oxygen density. In this case, the burning
reaction is substantially completed at the upstream side
surface of the catalyst layer 8, and the surface temper-
ature reaches about 860°C. At this instant, the
auxiliary catalyst layer 13 is heated only by the exhaust
25 gas discharged from the catalyst layer 8, and the surface
temperature thereof is as low as about 550°C. Even when
the oxygen density is further lowered, the temperature
difference between the catalyst layer 8 and the auxiliary

1 catalyst layer 13 is maintained almost constant,
because the oxygen amount is still sufficient (the actual
oxygen excessive rate is about 1.3 to 1.4 in the case
where the oxygen density becomes 15%). If the air amount
5 to be supplied to the mixing room 4 is decreased by about
30%, the air ratio at the catalyst layer 8 become 1.3 to
1.4. In this condition, for obtaining a perfect combus-
tion, the oxygen density more than 20% is required, and
when the oxygen density become as low as 18%, the actual
10 oxygen excessive rate become 1.1 to 1.2, thereby causing
a fear to produce carbon monoxide or unburned gas.
These combustible compositions are mixed with the air
supplied from the secondary air port 17 and flowed toward
the auxiliary catalyst layer 13, where a burning reaction
15 is effected. As a result, at the catalyst layer 8, the
burning reaction becomes weaker and the temperature
becomes lower, while at the auxiliary catalyst layer 13,
the burning reaction becomes stronger and the temperature
becomes higher. When the oxygen density is furthermore
20 lowered, the burning reaction becomes further weaker at
the catalyst layer 8 and further stronger at the auxiliary
catalyst layer 13. As a result, the temperatures of
these two layers gradually approach to each other, and
finally will be reversed. Now, by presetting a suitable
25 temperature difference value and controlling the pump 2
so as to stop the fuel supply when the temperature dif-
ference becomes lower than the preset value, the burning
in an oxygen starvation state can be prevented, and

1 the adverse influence on human being and beasts can be
avoided.

Requirement for setting the temperature difference depends on the target value of the oxygen limit density, the total amount of the burning, the area ratio of the catalyst layer 8 to the catalyst layer 13, and the predetermined air ratio, and it may be set in the control circuit 12. A suitable action can be easily carried out in response to a change of the total burning amount, if the predetermined temperature difference is previously stored in the control circuit 12. If the air supply rate to the mixing room 4 is maintained at the above-mentioned limit value, the operation may be apt to become unstable when the fuel supply amount or 15 the air supply amount changes. For effecting a perfect combustion at the catalyst layer 8, it is basically preferred to supply sufficient air. Therefore, it is suitable to practise the above-mentioned air flow change process only for a short time such as 2 to 3 minutes at 20 constant intervals of such as 30 minutes or one hour.

Fig. 6 shows a fifth embodiment, where there is provided a flow controller 18 including an opening and closing valve located at the middle of the secondary air tube 16 for opening the flow tube for a short time 25 at certain intervals. When the flow controller 18 is opened, a part of the air to be supplied to the mixing room 4 is supplied to the secondary air port 17 through the secondary air tube 16. As a result, the air supplied

1 to the mixing room 4 is decreased, and at the same time,
an air supply to the upstream side of the auxiliary
catalyst layer 13 is started, thereby producing the same
effects as in the fourth embodiment. In this embodiment,
5 no special operation of the fan 3 is required, and
since no excessive air is supplied from the secondary air
port 17 in a normal burning operation, the auxiliary
catalyst layer 13 is not cooled, and can be maintained
at a sufficiently high temperature, thereby assuring a
10 perfect purifying power against unburned composition or
carbon monoxide.

Next, a sixth embodiment will be described.

In the arrangement shown in Fig. 3, platinum (Pt) is carried by the catalyst layer 8, and a composition
15 produced by mixing palladium (Pd) and platinum at a weight ratio 2 : 1 is carried by the catalyst layer 13. The thickness of the catalyst layer 13 is about 80% of that of the catalyst layer 8, and the area of the former is about 30% of that of the latter, and the external
20 volume of the former is about 24% of that of the latter. The cell density (number of the communicating holes 8a, 13a per unit area) of the honeycomb which constitutes the carrier is 300 cells/in²) regarding the catalyst layer 8, while 400 cells/in² regarding the catalyst layer 13,
25 and accordingly, the diameter of the communicating holes 8a is smaller than that of the communicating holes 13a by about 30%.

As mentioned above, the catalyst layer 8 and

1 the catalyst layer 13 carry different precious metals,
and there is also a difference between the reacting
features of Pt and Pd on CO and kerosine as shown in
Fig. 7. Namely, Pd has a higher activity in oxidizing
5 of CO (here, 400 ppm CO is contained in the air), and
in particular, a superior activity at low temperature.
on the other hand, Pt has a higher activity in oxidizing
of kerosine (here, 2% kerosine vapor is contained in the
air), and has a perfect reacting feature (activity at
10 a condition of near 100% transforming rate) which is
significantly different from that of Pd. Therefore, in
the arrangement of Fig. 3, Pt is used at the catalyst
layer 8 for obtaining a superior burning reaction with
kerosine, while Pd is mainly used at the auxiliary
15 catalyst layer 13, which has a low temperature, for
purifying CO, which constitutes a main reactive composition,
efficiently at a low temperature. Although the reaction
starting feature at the catalyst layer 8 is expected to be
improved by mixing Pd, it is desired, for making the burn-
20 ing reaction more perfect, to use Pt only or Pt as a
main composition. On the other hand, at the auxiliary
catalyst layer 13, although Pd only may be used for
purifying CO, Pt is preferred to be mixed in consideration
of the fuel slip due to the activity deterioration or
25 locally lowered temperature of the catalyst layer 8. With
respect to the reactivity on the fuel, the above-mentioned
activity difference is seen in gaseous fuels such as
propane or butane similarly to the above-mentioned kerosine,

1 and any gaseous fuel excluding methane has the same
feature.

Even if the volume of the auxiliary catalyst
layer 13 is equal to that of the catalyst layer 8, there
5 is no problem with respect to the performance. However,
since a great size of the auxiliary catalyst layer 13
causes a high cost, an excessive size thereof is
undesirable in the practical view point. The load on the
auxiliary catalyst layer 13 is usually small, and a perfect
10 reaction can be obtained, even if the spacial speed is
considerably increased. Fig. 8 shows a relation between
the volume ratio of the auxiliary catalyst layer 13 to the
catalyst layer 8 and the transforming rate of the reactive
substances. In an initial stage where the CO density is
15 below 100 ppm, a perfect purification can be obtained,
even when the volume ratio of the auxiliary catalyst
layer 13 to the catalyst layer 8 is made as low as 10%
and the spacial gass speed is increased by about ten
times. Even in a condition where no reaction is caused
20 at the catalyst layer 8 (all fuel slips and reaches the
auxiliary catalyst layer 13), an almost normal burning
can be effected if the volume ratio of the auxiliary
catalyst layer 13 is as great as 50%, thereby preventing
a great amount of smell or CO from being exhausted, and
25 preventing any abnormal condition such as a back-fire.
An abnormality of the catalyst layer 8 can be detected
by measuring the temperature rise of the auxiliary
catalyst layer 13 by means of the thermocouple 14, and

1 in response to this detected abnormality, the burning
can be stopped. In consequence, considering the cost
requirement, it is required to make the size of the
auxiliary catalyst layer 13 minimum, and therefore, the
5 volume ratio of the auxiliary catalyst layer 13 to the
catalyst layer 8 may be preferably selected at 10 to 50%
according to the precision of the temperature detection
and the allowable value for deterioration of the catalyst
layer 8.

10 The density of the unburned composition passing
through the auxiliary catalyst layer 13 is far thin in
comparison with that through the catalyst layer 8, and
as a result, the diffusion of the reactive substance for
oxidizing reaction become important. If the diameter
15 of the communicating holes 13a of the auxiliary catalyst
layer 13 is made smaller, in other words, the honeycomb
cell density is made greater, the diffusion time of
the unburned composition can be shortened and the
reactivity is improved, resulting in a high transforming
20 rate even at a low temperature, as shown in Fig. 9. In
case of the catalyst layer 8, excessive cell density
causes a reaction heat concentration and an excessive
temperature rise, thereby deteriorating the catalytic
activity. In case of the auxiliary catalyst layer 13,
25 however, there is no such deterioration, because the
produced heat is small due to the thin density of the gas.
Fig. 9 indicates that if the cell density is increased,
the reactivity is improved and the purification becomes

1 perfect, even in case the volume of the auxiliary catalyst
layer 13 is small (spacial speed is great). This structure
is helpful for decreasing the size of the auxiliary catalyst
layer 13 through which a gas of low temperature and low
5 density passes. The greater density of the cell is
accompanied with an increased flow resistance, and the cell
density has an upper limit due to the restriction in
fabrication. However, by making the diameter of the com-
municating holes 13a of the auxiliary catalyst layer 13
10 smaller than that of the communicating holes 8a of the
catalyst layer 8, it become possible to purify the exhaust
gas efficiently with a small volume and with a low cost.

In every case mentioned above, the carrier
of the catalyst layer 8 or the auxiliary catalyst layer
15 13 is not limited to a ceramic honeycomb as shown in
the above-mentioned embodiments, but a ceramic foam,
a braided body of anti-heat fibers, or a metal honeycomb
can be used with the same advantage obtained. The above-
mentioned advantage is not influenced by the kind or the
20 shape of the carrying body of the catalyst layer 8 or
the auxiliary catalyst layer 13.

INDUSTRIAL APPLICABILITY

As mentioned above, in a catalytic burning
apparatus according to the present invention, an uniform
25 catalyst preheating can be effected in a short time,
because the catalyst layer is preheated by utilizing a
flame burning which produces an hot exhaust gas. Further,

1 since it is confirmed by means of ion current detecting
means that a stable flame is formed in a flame burning
stage, and no flame is formed in a catalytic burning
stage, any effusion of unburned gas due to misignition
5 or misfire can be prevented. In addition, in a catalytic
burning, it can be confirmed that there is not any
backfire phenomenon, which may be caused by an overheating
of the catalyst layer due to an abnormality of the pump
or the fan and may form a flame at the flame ports.

10 Further, by providing temperature detecting means for
the catalyst layer, the preheat temperature of the
catalyst layer can be suitably adjusted and a catalytic
burning realizing a perfect reaction can be started
from the initial stage. In case of an abnormal structure
15 or an abnormal activity of the catalyst layer, the
abnormality can be quickly detected and any smell or carbon
monoxide due to an imperfect combustion can be prevented
from being produced. By conducting flame burnings at
certain intervals and confirming by ion electric current
20 detecting means that a predetermined electric current
is flowing, any abnormality of the oxygen density can be
detected, and any oxygen starvation having a harmful
influence on the human body can be prevented. By providing
two stages of catalyst layers and detecting the tempera-
25 ture difference between these two catalyst layers, any
activity deterioration or damage of the catalyst layers
can be detected, and further, by supplying a secondary air
to the upstream side of the catalyst layer (auxiliary

1 catalyst layer) located at the downstream side, any
oxygen starvation can be detected. By using Pt as a
main composition for the upstream side catalyst layer,
and Pd as a main composition for the downstream side
5 catalyst layer, an optimum reaction suitable to the
composition to be burned or the density of the same can
be effected, thereby providing a burning apparatus
capable of effecting a perfect reaction. By making smaller
the volume of the downstream side catalyst layer having a
10 smaller load, or making smaller the cell diameter of the
downstream side catalyst layer having a lower combustible
gas density, an efficient burning and an efficient exhaust
gas purification can be effected at low cost.

CLAIMS

1. A catalytic burning apparatus including a mixing room for mixing fuel with air, flame ports arranged downstream of said mixing room, a catalyst layer disposed downstream of said flame ports and bored with a plurality of communication holes, and ion current detecting means and igniting means disposed near said flame ports, the arrangement being such that the igniting means is operated for forming a flame at the flame ports, the flame is extinguished after a predetermined time length by once stopping the fuel supply, and then a burning reaction on the surface of the catalyst layer is started by supplying fuel again without operating the igniting means, characterized in that:

the ion current detecting means detect the condition that a predetermined electric current value is not obtained and the burning is controlled to be stopped when the flame is formed at the flame ports, and the ion current detecting means detect the condition that a predetermined electric current value is obtained and the burning is controlled to be stopped when the burning reaction is started at the catalyst layer.

2. A catalytic burning apparatus claimed in claim 1, wherein the apparatus further comprises temperature detecting means for detecting the temperature of the catalyst layer, and the flame burning time is controlled such that, after the flame forming at the flame ports, when the temperature of the catalyst layer

reaches a predetermined value, the step is shifted to the step of temporarily stopping the fuel supply.

3. A catalytic burning apparatus claimed in claim 1, further comprising control means which operates the igniting means at predetermined intervals for forming a flame at the flame ports for a predetermined time, stops the fuel supply when the ion current detecting means detects the condition that the predetermined electric current is not obtained, and restarts the catalytic burning through steps of temporary stopping of the fuel supply and resupplying of the fuel, when the ion current detecting means detect condition that the predetermined electric current is obtained.

4. A catalytic burning apparatus claimed in claim 1, further comprising

an auxiliary catalyst layer arranged downstream of the catalyst layer and bored with a plurality of communicating holes,

temperature detecting means for detecting temperatures of said catalyst layer and said auxiliary catalyst layer,

a secondary air supply section having an opening at the upstream side of the auxiliary catalyst layer,

control means for decreasing the air supply to the mixing room by a predetermined ratio at predetermined intervals, and

control means interconnected with said temperature

detecting means for stopping the fuel supply, when the temperature difference between these two catalyst layers becomes below a predetermined value.

5. A catalytic burning apparatus claimed in claim 4, further comprising,

air supply means communicating with both of said mixing room and said secondary air supply section,

flow control means for making a communication with said secondary air supply section at predetermined intervals for a predetermined time, and

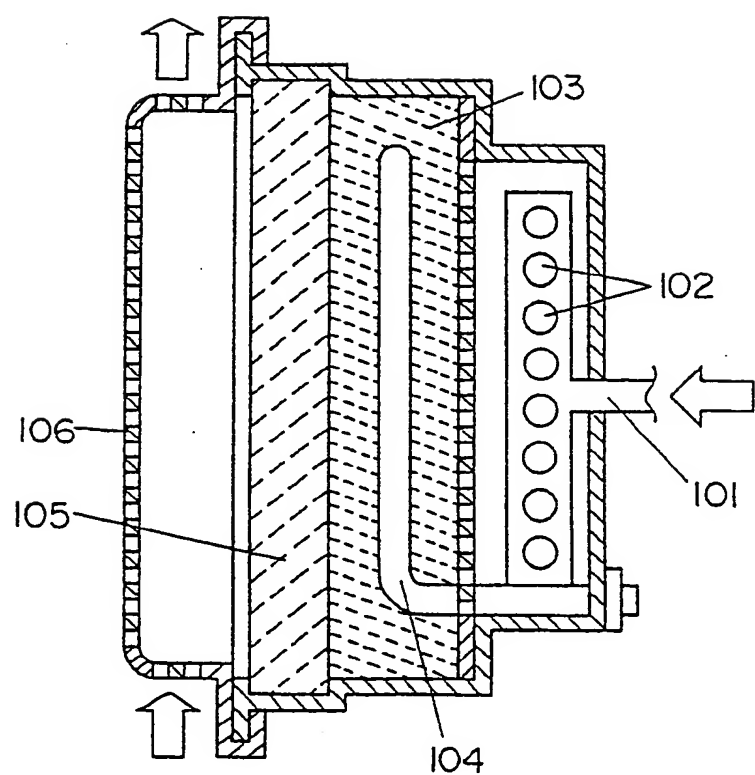
control means interconnected with said temperature detecting means for stopping the fuel supply, when the temperature difference between these two catalyst layers become below a predetermined value.

6. A catalytic burning apparatus claimed in claim 1 or 4, wherein the apparatus comprises an auxiliary catalyst layer arranged downstream of the catalyst layer and bored with a plurality of communicating holes, and said catalyst layer carries platinum or a mixed precious metal mainly composed of platinum, while said auxiliary catalyst layer carries palladium or a mixed precious metal mainly composed of palladium.

7. A catalytic burning apparatus claimed in any one of claims 1, 4 and 6, wherein the apparatus comprises an auxiliary catalyst layer arranged downstream of the catalyst layer and bored with a plurality of communicating holes, and the volume of the auxiliary catalyst layer is 10 to 50% of that of the first-mentioned catalyst layer.

8. A catalyst burning apparatus claimed in one of claims 1, 4, 6 and 7, wherein the apparatus comprises an auxiliary catalyst layer arranged downstream of the catalyst layer and bored with a plurality of communicating holes, and the diameter of the communicating holes of the auxiliary catalyst layer is smaller than that of the first-mentioned catalyst layer.

FIG. 1



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FIG. 2

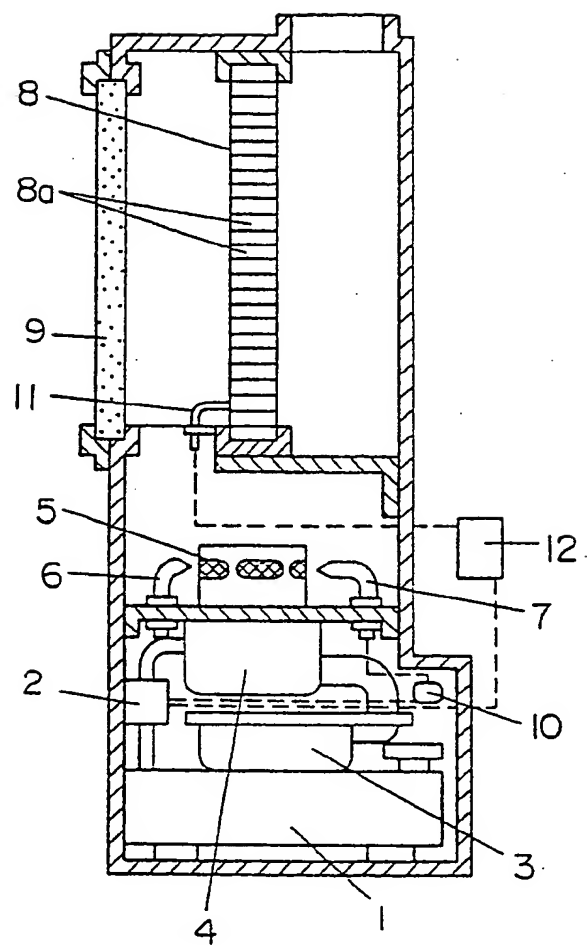


FIG. 3

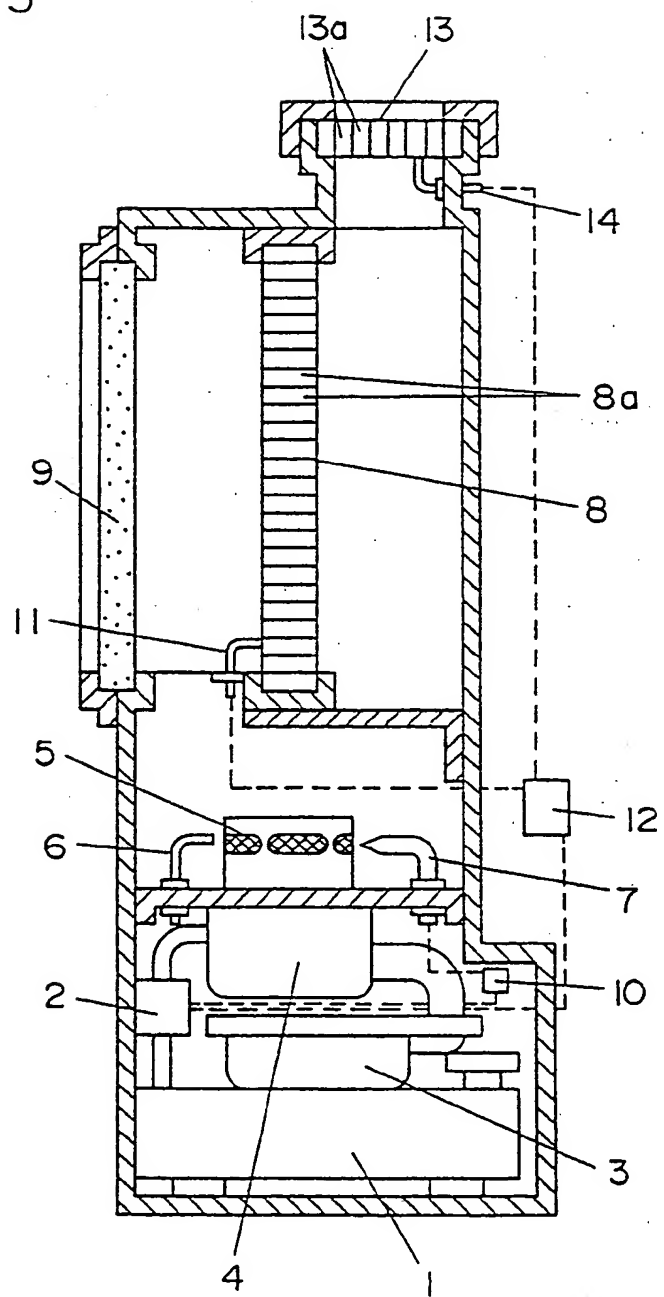
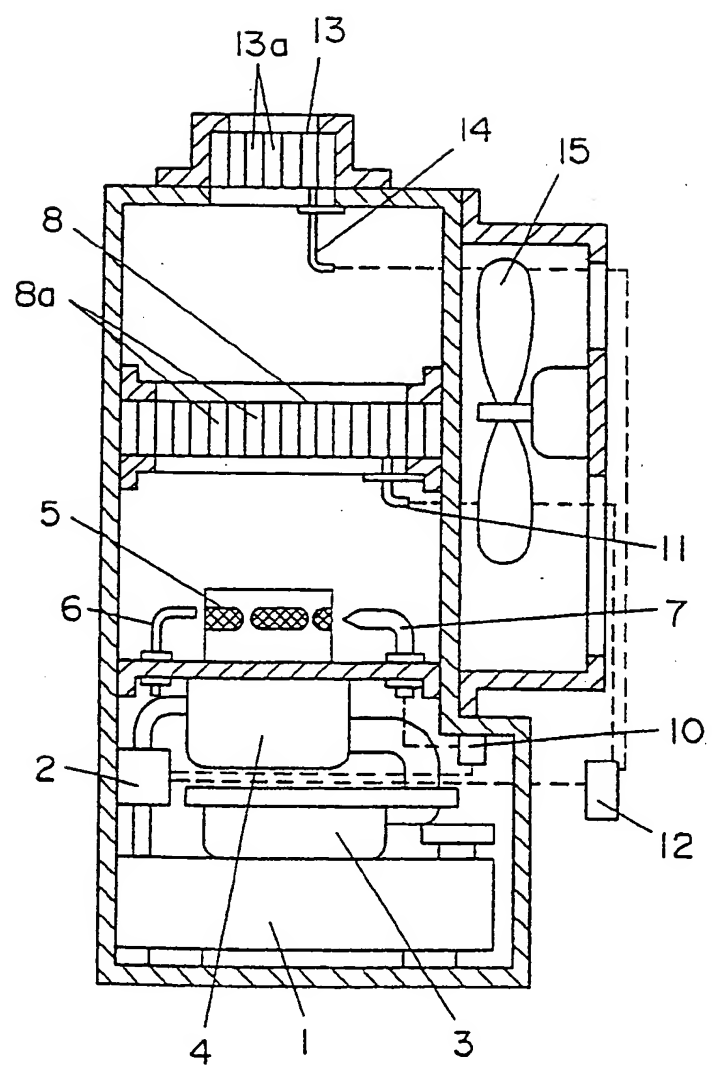
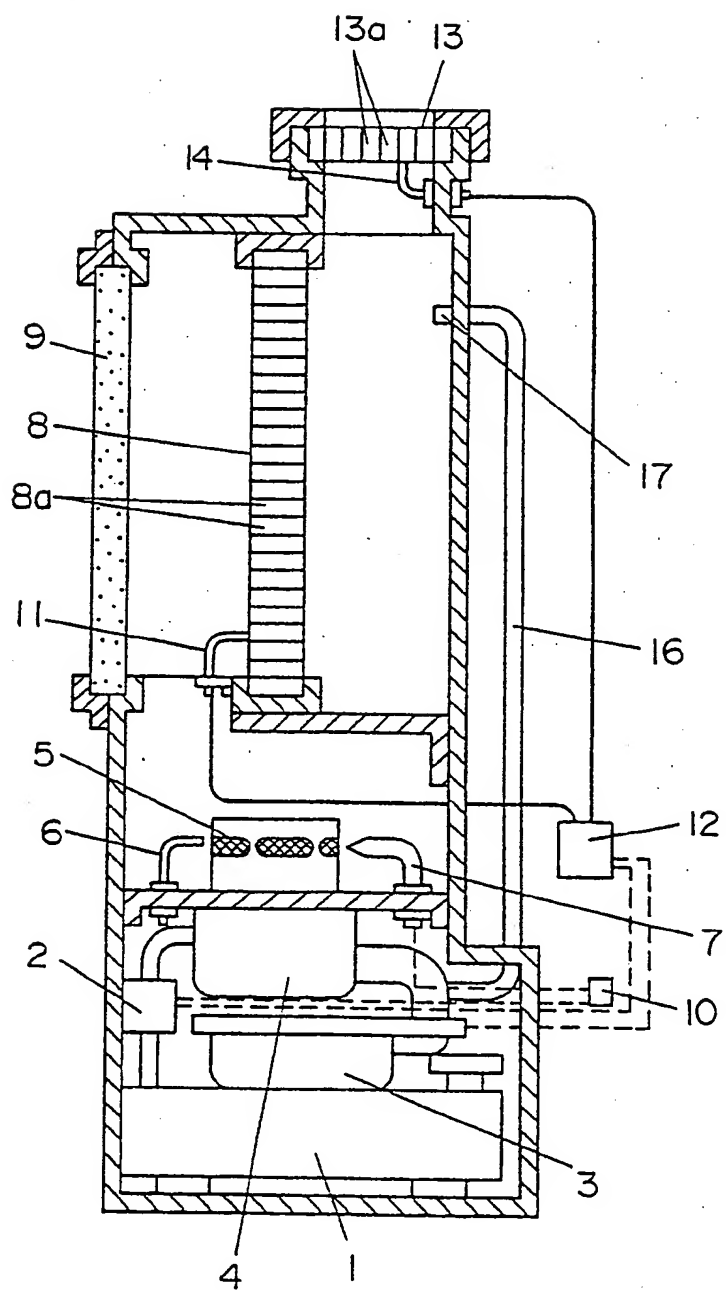


FIG. 4



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FIG. 5



220 000 000

FIG. 6

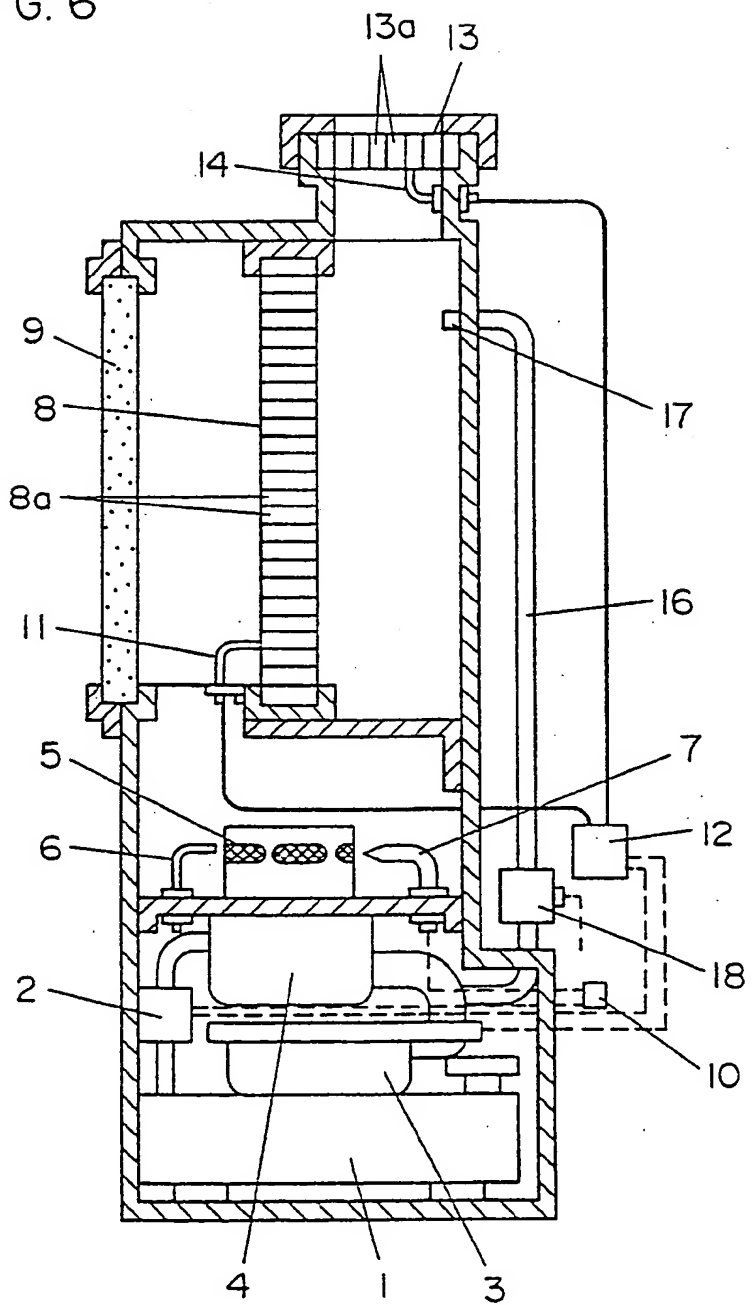


FIG. 7

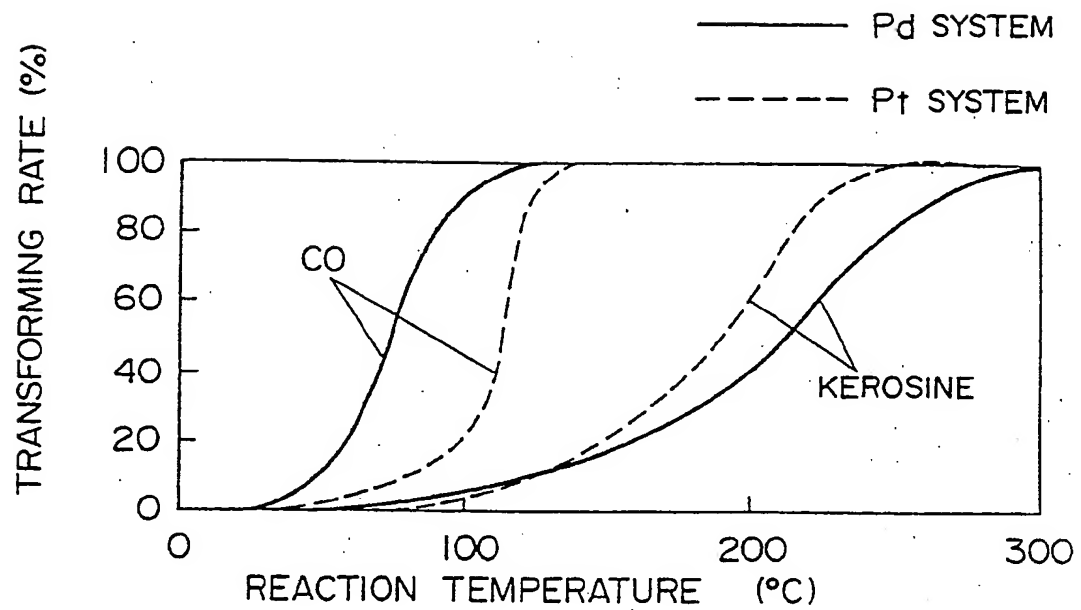
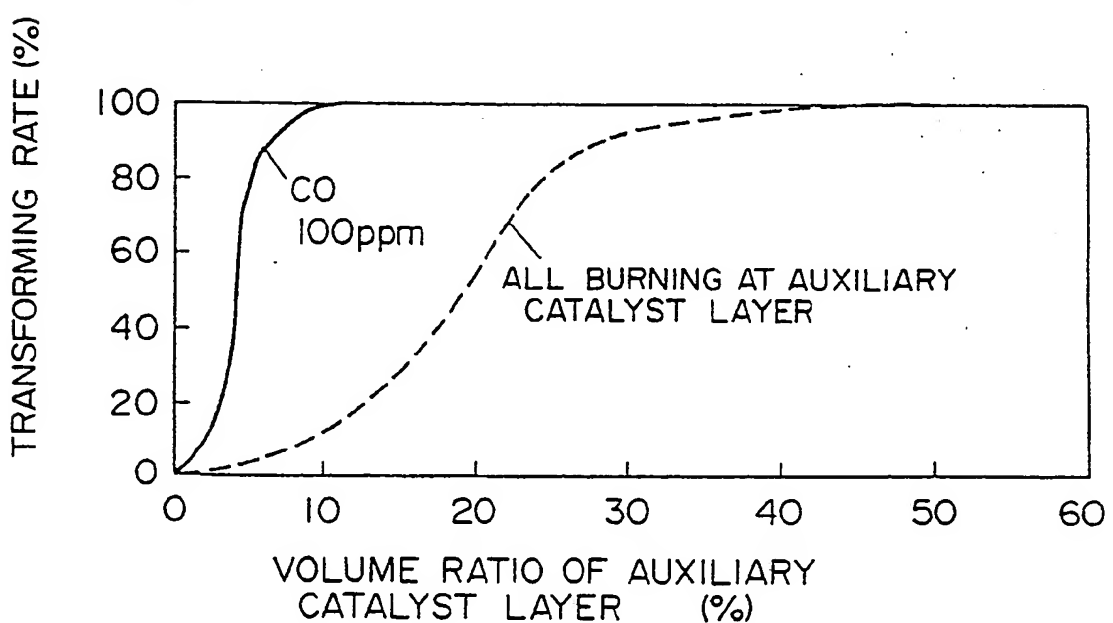
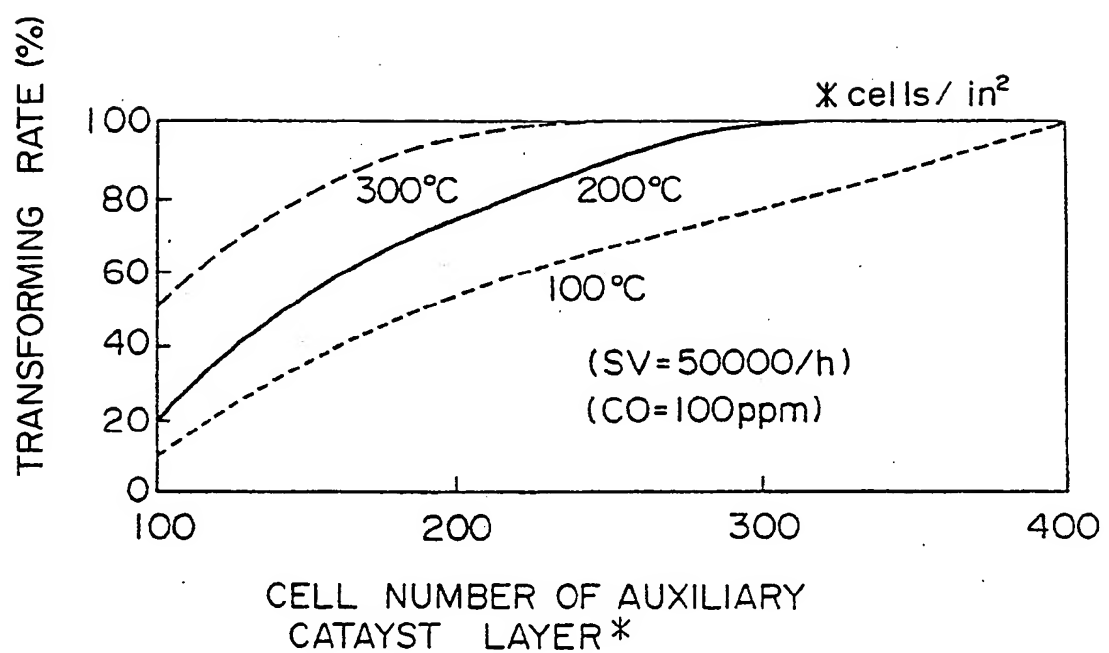


FIG. 8



22 22 22 22 22
22 22 22 22 22

FIG. 9



LIST OF THE NUMBERED MEMBERS IN THE DRAWINGS

- 1 --- Tank,
- 2 --- Pump,
- 3 --- Fan,
- 4 --- Mixing room,
- 5 --- Flame port,
- 6 --- Ignition plug,
- 7 --- Flame rod,
- 8 --- Catalyst layer,
- 8a, 13a --- Communicating hole,
- 9 --- Transparent window,
- 10 --- Controller,
- 11, 14 --- Thermocouple,
- 12 --- Control circuit,
- 13 --- Auxiliary catalyst layer,
- 15 --- Air blast fan,
- 16 --- Secondary air tube,
- 17 --- Secondary air port,
- 18 --- Flow controller,
- 101 --- Fuel tube,
- 102 --- Injection port,
- 103 --- Insulator,
- 104 --- Electric heater,
- 105 --- Catalyst layer,
- 106 --- Cover.

INTERNATIONAL SEARCH REPORT

International Application No. PCT/JP89/00795

I. CLASSIFICATION OF SUBJECT MATTER (If several classification symbols apply, indicate all) *

According to International Patent Classification (IPC) or to both National Classification and IPC

Int. Cl⁴ F23N5/12, F23N5/02, F23D14/18, F23C11/00

II. FIELDS SEARCHED

Minimum Documentation Searched :

| Classification System : | Classification Symbols |
|-------------------------|--|
| IPC | F23N5/12, F23N5/02, F23D14/18, F23C11/00, F23N5/10 |

Documentation Searched other than Minimum Documentation
to the Extent that such Documents are Included in the Fields Searched *

| | |
|---------------------------|-------------|
| Jitsuyo Shinan Koho | 1926 - 1989 |
| Kokai Jitsuyo Shinan Koho | 1971 - 1989 |

III. DOCUMENTS CONSIDERED TO BE RELEVANT *

| Category * | Citation of Document, if with indication, where appropriate, of the relevant passages :- | Relevant to Claim No. |
|------------|--|-----------------------|
| Y | JP, A, 59-13821 (Mitsubishi Electric Corporation) 24 January 1984 (24. 01. 84) Column 1, line 16 to column 3, line 10, Fig. 1 (Family : none) | 1 - 8 |

| | | |
|---|---|---|
| Y | JP, A, 62-162821 (Matsushita Electric Ind. Co., Ltd.) 18 July 1987 (18. 07. 87) Column 3, lines 11 to 20, column 5, lines 3 to 14, Fig. 1 (Family : none) | 4 |
|---|---|---|

| | | |
|---|--|------|
| Y | JP, A, 62-41511 (Nippon Shokubai Kagaku Kogyo Co., Ltd.) 23 February 1987 (23. 02. 87) Column 3, lines 2 to 8 (Family : none) | 4, 6 |
|---|--|------|

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"Z" document member of the same patent family

IV. CERTIFICATION

| | |
|---|---|
| Date of the Actual Completion of the International Search | Date of Mailing of this International Search Report |
|---|---|

| | |
|-------------------------------|-------------------------------|
| October 27, 1989 (27. 10. 89) | November 6, 1989 (06. 11. 89) |
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International Searching Authority

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